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Cancer Discovery at One Year: The Editors’ Interim Analysis. vi
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RESEARCH BRIEFS
ATM Mutations in Patients with Hereditary Pancreatic Cancer. 41
Précis: Next-generation sequencing identifies inherited ATM mutations in kindreds with hereditary pancreatic ductal adenocarcinoma.

Molecular Ontogeny of Donor-Derived Follicular Lymphomas Occurring after Hematopoietic Cell Transplantation. 47
Précis: Analysis of a donor–recipient pair with follicular lymphoma reveals the time-course of somatic mutations acquired during lymphomagenesis.
RESEARCH ARTICLES

Genomic Complexity and AKT Dependence in Serous Ovarian Cancer .................56

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High-Throughput Detection of Actionable Genomic Alterations in Clinical Tumor Samples by Targeted, Massively Parallel Sequencing .........................82

Précis: Targeted, sequencing-based profiling of archival tumor samples identifies genetic alterations that can direct personalized therapy.

Loss of the 14-3-3σ Tumor Suppressor Is a Critical Event in ErbB2-Mediated Tumor Progression .........................68
C. Ling, V-M-T. Su, D. Zuo, and W.J. Muller

Précis: 14-3-3σ inactivation accelerates formation and promotes metastasis of ErbB2/HER2-induced tumors.

For more News and Research Watch, visit Cancer Discovery online at www.AACR.org/CDnews. Online-only News stories include the following:

• Biotech Firms Look for Virtual Success  • Dual HER2 Blockade Slows Metastatic Breast Cancer
• “Reversed” Krebs Cycle Can Feed Tumors  • Modified Stem Cells Create Tumor-Attacking T Cells

ON THE COVER

Wagle and colleagues describe a method to profile clinically relevant mutations in formalin-fixed, paraffin-embedded tumor samples involving exon capture of frequently mutated or polymorphic genes followed by massively parallel sequencing. This method identifies single-nucleotide variants, insertions, deletions, and copy number alterations overlooked by current genotyping-based methods with high specificity and sensitivity. Identification of such “actionable” genetic alterations that predict response to targeted or conventional cytotoxic therapies has the potential to facilitate individualized cancer treatment in a time- and cost-effective manner. For details, please see the article by Wagle and colleagues on page 82.